

A Further Investigation of the
Chlorides of
Paranitroorthosulphobenzoic Acid

DISSERTATION

SUBMITTED TO THE BOARD OF UNIVERSITY STUDIES OF THE
JOHNS HOPKINS UNIVERSITY FOR THE DEGREE
OF DOCTOR OF PHILOSOPHY

—BY—

FREDERICK S. HOLLIS

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I. INTRODUCTION.

The present investigation may be divided into two parts. The first consists of further work on the method of preparation and separation of the chlorides of paranitroorthosulphobenzoic acid. This work was confined largely to the unsymmetrical or low-melting chloride, as this is the one used mainly in the second part of the investigation. The preparation of the unsymmetrical dichloride, first obtained by Gray,¹ was found to be a matter of considerable difficulty and uncertainty, unless the crystallization could be conducted out of doors at a very low temperature.

As the result of a series of experiments, undertaken to determine the best conditions for the preparation of the unsymmetrical chloride, a method has been adopted by which the unsymmetrical chloride may be prepared in any desired quantity in the laboratory.

The second part of the investigation consists of a study of the action of benzene and aluminium chloride on the chlorides under varying conditions, and the preparation of a series of derivatives of the product formed.

Saunders² and McKee,³ working with the chlorides of orthosulphobenzoic acid, found that the action of benzene and aluminium chloride gives the same products with both chlorides.

It was thought that, on account of the greater stability of the paranitroorthosulphobenzoic acid, the action of benzene and aluminium chloride might lead to the formation of two series of derivatives, one derived from the symmetrical chloride and the other from the unsymmetrical chloride, in which the resulting compound would retain the unsymmetrical structure of the chloride. This proved not to be the case, as the product of the action of benzene and aluminium chloride on both of the chlorides was found to be paranitroorthobenzoyl-benzenesulphone chloride.

¹ Inaug. Diss., Johns Hopkins University, 1895.

² Am. Chem. J., 17, 355.

³ Inaug. Diss. Johns Hopkins University, 1895.

It was found that paranitroorthosulphobenzoic acid does not yield a sulphone corresponding to the one obtained from orthosulphobenzoic acid by Saunders,¹ although the reaction was conducted under the conditions used by him, as well as under a variety of different conditions.

A series of derivatives of paranitroorthobenzoylbenzenesulphone chloride was prepared.

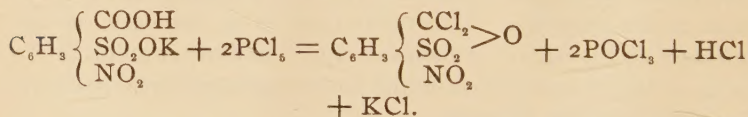
II. Preparation of Material.

The acid potassium salt of paranitroorthosulphobenzoic acid was prepared from paranitrotoluene according to the method described by Kastle,² and used later by Gray.³ This consists in introducing the sulphonic acid group into paranitrotoluene and passing through the calcium salt to the potassium salt of paranitrotolueneorthosulphonic acid. The potassium salt is oxidized by means of potassium permanganate to the potassium salt of paranitroorthosulphobenzoic acid, and this is converted into the acid potassium salt by the action of hydrochloric acid.

One thousand grams of paranitrotoluene gave 1439 grams of the neutral potassium salt of paranitrotolueneorthosulphonic acid. One thousand grams of the potassium salt of paranitrotolueneorthosulphonic acid gave 800 grams of the acid potassium salt of paranitroorthosulphobenzoic acid.

III. The Action of Phosphorus Pentachloride on the Acid Potassium Salt of Paranitroorthosulphobenzoic Acid.

The action of phosphorus pentachloride on the anhydrous acid potassium salt of paranitroorthosulphobenzoic acid gives rise to the formation of an unsymmetrical and a symmetrical dichloride, as determined by Gray,⁴ according to the following equations :

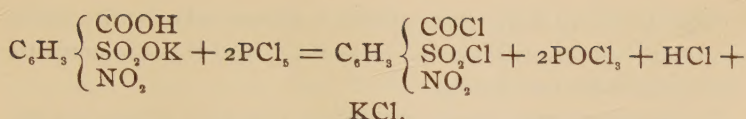


¹ Am. Chem. J., **17**, 363.

² Am. Chem. J., **11**, 171.

³ Inaug. Diss. Johns Hopkins University, 1895.

⁴ Inaug. Diss. Johns Hopkins University, 1895.



Gray¹ found that the relative amount of each chloride depends on the temperature and length of time which the phosphorus pentachloride is allowed to act on the acid potassium salt. The largest amount of the unsymmetrical chloride, amounting to 80 or 90 per cent. of the total, was formed by heating a mixture of 2 molecules of phosphorus pentachloride and 1 molecule of the anhydrous acid potassium salt to 150° C. for four or five hours in a distilling bulb, immersed in a sulphuric-acid bath.

Under these conditions, the amount of symmetrical chloride formed amounted to 10 or 20 per cent. of the total, but this is increased to 30 per cent. by heating in an open dish on a water-bath. The symmetrical chloride was separated by using chloroform as a solvent.

As considerable difficulty was experienced, mainly in the preparation of the unsymmetrical chloride, according to the directions given by Gray, a series of experiments was made under different conditions with amounts of the acid potassium salt varying from 20 to 60 grams in order to determine the conditions most favorable for the formation of the unsymmetrical chloride. The results of these experiments are embodied in the following method :

a. The Preparation of the Unsymmetrical Chloride.—A mixture of 1 molecule of the acid potassium salt, previously heated to 150° C. for four hours, and 2½ molecules of phosphorus pentachloride is carefully ground together in a mortar and introduced into a distilling bulb having a capacity of about six times that of the volume of the mixture. The outlet tube of the bulb is closed, and a cork, through which runs a glass tube about three feet long, the lower end reaching nearly to the surface of the mixture, inserted in the neck.

¹ Inaug. Diss., Johns Hopkins University, 1895.

The distilling bulb, thus closed, is immersed in a sulphuric-acid bath, previously heated to $150^{\circ}\text{C}.$, and this temperature maintained for five hours.

No reaction takes place on mixing the acid potassium salt and the phosphorus pentachloride, but, upon immersing the bulb containing the mixture in the heated bath, vigorous action begins immediately, and the resulting phosphorus oxychloride is conducted back upon the product by means of the condensing tube. The temperature of $150^{\circ}\text{C}.$ cannot safely be exceeded, as decomposition of the chloride begins at $160^{\circ}\text{C}.$

At the end of five hours the tube is removed from the neck of the flask, the perforated stopper replaced by a solid one, the outlet tube opened and the phosphorus oxychloride distilled off.

The resulting chloride, which is in the form of a thick, yellow, oily liquid, is poured into a large bottle, nearly filled with cold water, and shaken vigorously so as to break it into small globules. This washing is continued until the chloride hardens to a solid cake, which commonly takes place after washing with five or six successive portions of cold water. The solidified chloride is broken up and dried by pressing between filter paper, and in this form it may be exposed to the air without undergoing much, if any, decomposition.

By using two and a half molecules of phosphorus pentachloride, all of the acid potassium salt is converted into the form of the chloride, while, if but two molecules are used, varying amounts of the acid potassium salt, in some cases as much as 25 per cent., are unacted upon and may be recovered from the wash-water.

The product obtained by this method consists, with the exception of slight impurities, of only the unsymmetrical chloride, thus making the crystallization from chloroform unnecessary. One hundred grams of anhydrous acid potassium salt give, on an average, 99 grams of the crude unsymmetrical chloride.

Previous work showed that the crystallization of the un-

symmetrical chloride from the ligroïn (50–80°) was a difficult matter, unless it could be done out of doors during very cold weather. The same was found to be the case with the unsymmetrical chloride of orthosulphobenzoic acid by Saunders.¹

A solution of the crude chloride in purified ligroïn (50–80°), on standing out of doors at a temperature of 6° F., crystallized in clusters of crystals, some of which measured three centimeters in length. An attempt was made to obtain the chloride in the form of crystals by cooling the ligroïn solution to 0° C. in a refrigerating box, such as was used by McKee.² The chloride invariably separated as an oil, and no advantage was derived by drawing a current of cold, dry air through the flask containing the ligroïn solution. Some of the oily chloride which separated from the ligroïn solution, and from which the ligroïn had been decanted, formed an opaque semi-solid mass on placing it in a freezing mixture, but no crystals were deposited.

The first indication of crystallization in the laboratory was obtained on placing some of the oil, which had been dissolved several times in fresh portions of hot ligroïn (50–80°) and allowed to separate out on cooling, in a freezing mixture and stirring with a rod. This suggested the possibility that some material, which by its presence retards crystallization, had been dissolved out of the mass by the several portions of ligroïn in which it had been dissolved. This was shown to be the case by several tests and gave rise to the following method of purification and crystallization, in which the further change is made of using ligroïn of boiling-point 90–125° as the solvent. This ligroïn is to be preferred to that having a lower boiling-point, as the chloride is apparently much more soluble in it and crystallizes from the solution at the temperature of the laboratory. The ligroïn is purified by shaking in a separating funnel with concentrated sulphuric acid until it imparts no color to a fresh portion of acid, after which it is treated with caustic soda solution to neutralize the acid, and washed free from alkali by water.

¹ Am. Chem. J., 17, 348.

² Inaug. Diss., Johns Hopkins University, 1895.

The crude chloride, which is always somewhat dark colored and gummy, is placed in an Erlenmeyer flask with purified ligroïn (50–80°) and washed by stirring it down with a rod until it yields a granular powder having but little color.

On boiling the chloride thus purified with ligroïn (90–125°), it dissolves, with the exception of slight remaining impurities, and on cooling the solution becomes cloudy and the excess of chloride separates out as a light-colored oil. It is best to decant the solution from the separated oil after separation has mainly ceased at the temperature at which crystallization is to proceed, but before the solution has become clear. The chloride is obtained from this solution at the temperature of the laboratory in clusters of needles having the form of long monoclinic prisms as observed by Gray.¹ The rate of crystallization is increased by keeping the solution at a lower temperature, but no especial advantage is derived unless the temperature is very low, when larger crystals are obtained. The crystals of the chloride thus obtained have a constant melting-point of 57° C. (uncor.).

The purified chloride which separates from the ligroïn as an oil generally crystallizes on standing, but it is better to redissolve it in a fresh portion of ligroïn, by which it is further purified, and proceed according to the directions given above.

The principal impurity which causes difficulty in the crystallization of the unsymmetrical chloride seems to be that which is removed by the preliminary treatment with ligroïn (50–80°).

On evaporating the ligroïn used in washing the chloride, this impurity remains as a dark-colored, slightly viscous liquid having a strong acid reaction. It showed no tendency to crystallize after standing in the laboratory for five months. A small amount of another impurity was obtained as a flocculent material on dissolving the crude chloride in chloroform. It melts after purification at 100–105° C., and is probably the anhydride.

¹ Inaug. Diss., Johns Hopkins University, 1895.

b. The Preparation of the Symmetrical Chloride.—One portion of the symmetrical chloride was made in order to test the action of benzene and aluminium chloride, but no comparative tests were made, as in the case of the unsymmetrical chloride. The method used for its preparation differed somewhat from that of Gray.

The conditions chosen for its preparation were, as far as possible, the opposite of those found to be most favorable for the formation of the unsymmetrical chloride.

The anhydrous acid potassium salt and phosphorus pentachloride, in the proportion of 1 molecule of the former to 2 molecules of the latter, are mixed by grinding them together, and the action commenced by placing the vessel containing the mixture in a sulphuric-acid bath previously heated to 150° C.

The vessel is removed from the bath as soon as the action commences, and allowed to stand until the action is complete, which requires about ten minutes.

The mass is washed, as in the case of the unsymmetrical chloride, by shaking in a bottle with five or six portions of cold water.

The chloride thus washed exists in the form of a light colored, thick gum. It is dissolved in anhydrous chloroform, the solution dried by means of calcium chloride, and allowed to stand at the temperature of the laboratory. Crystals of the symmetrical chloride are formed only after the chloroform has nearly all evaporated, and crystallization proceeds very slowly.

The chloride was obtained in the form of small monoclinic crystals, having a constant melting-point of 94° C. (uncor.).

The yield indicated that, under the conditions used, the action of the phosphorus pentachloride is incomplete, but between 35 and 40 per cent. of the product obtained consists of the symmetrical chloride.

IV. The Action of Benzene and Aluminum Chloride on the Chlorides of Paranitroorthosulphobenzoic Acid.

a. The Action on the Unsymmetrical Chloride.—When alumi-

nium chloride is added to a solution of the unsymmetrical chloride in benzene, slight action begins immediately, as is shown by the darkening of the color of the solution and a slight evolution of hydrochloric acid gas.

In the case of small quantities of the chloride, heated with an excess of aluminium chloride, the reaction is complete in from fifteen to twenty minutes. A series of experiments was made in order to determine the best conditions for conducting the reaction and for separating and purifying the product.

Under some conditions the product contains a considerable amount of a dark purple material which separates as an impurity on crystallization. As a result of these experiments, the following method was adopted:

Twenty grams of the unsymmetrical chloride is dissolved in 100 cc. of benzene in a flask provided with a return-condenser, and about ten grams of aluminium chloride in small pieces added. This is heated with a small flame for from one to two minutes, at the end of which time vigorous action commences and continues without further heating for ten minutes.

After the action is over, the flask is heated repeatedly so as to maintain an even evolution of hydrochloric acid gas for about eight minutes, at the end of which the reaction is complete. The resulting product is poured into 750 cc. of water in a liter separating-funnel, 75 cc. of hydrochloric acid (sp. gr. 1.12) added and the mixture well shaken.

After the benzene layer has risen, the water is drawn off, and the small amount of the product suspended in it separated by filtration. The greater part of the product is in the form of a pinkish-white powder, which remains in suspension in the benzene, from which it is separated by filtration, dried and purified with the portion obtained from the water. The product is purified by dissolving in benzene and adding rather more than an equal volume of anhydrous ether, which causes a more rapid crystallization. By this method it is obtained in clusters of small, apparently monoclinic crystals, having a rhombohedral habit, or as a granular powder if crystallization takes place rapidly.

Some of the larger crystals measured four or five millimeters on an edge. The larger crystals have a purple or green color, and the granular form is generally slightly green. Both forms yield a white powder. The pure crystalline product has a constant melting-point of 177°C . (uncor.).

The portion of the product which remains in solution in the benzene is mixed with a small amount of the purple impurity before described, but it is obtained in a crystalline form of fair purity by drying the benzene solution, evaporating it to about one-third of its volume, and adding rather more than an equal volume of absolute ether. The dark impurity may be largely removed by washing with absolute alcohol in which it dissolves readily.

An attempt was made to prevent the darkening of the benzene solution during evaporation by conducting the evaporation in a current of sulphur dioxide, as it was believed that the darkening was due, in part at least, to the action of the air.

The product was not materially improved and a certain amount of free acid was always found to be present after such treatment. The rapid evaporation of the dried benzene solution and the addition of an equal volume of absolute ether to insure rapid crystallization is greatly to be preferred.

Twenty grams of the unsymmetrical chloride gave sixteen grams of the product as first obtained. This was slightly decreased by recrystallization.

The product has a characteristic disagreeable odor.

b. The Action on the Symmetrical Chloride.—The reaction was conducted exactly as in the case of the unsymmetrical chloride, using the same relative quantities and method of treatment. The method of separation was also the same.

The product gave, on purification, crystals of the same form, size and color as those obtained from the unsymmetrical chloride.

The melting-point of the purest crystals is also the same, 177°C . (uncor.).

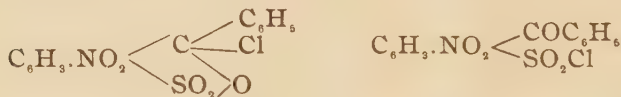
Two grams of the symmetrical chloride gave one and a half grams of the product.

Analyses of the product of the action of benzene and aluminum chloride on the unsymmetrical chloride :

0.2051	gram	gave	0.3632	gram	CO ₂ .
0.2992	"	"	0.5249	"	"
0.1994	"	"	0.0535	"	H ₂ O.
0.2992	"	"	0.0819	"	"
0.2508	"	"	9.03	cc.	N.
0.3496	"	"	12.88	"	"
0.2601	"	"	0.1957	gram	BaSO ₄ .
0.2058	"	"	0.1584	"	"
0.2602	"	"	0.1888	"	"
0.2492	"	"	0.1878	"	"
0.2424	"	"	0.1811	"	"
0.2434	"	"	0.1072	"	AgCl.
0.2058	"	"	0.0925	"	"
0.2602	"	"	0.1161	"	"

	Calculated for C ₁₃ H ₉ O ₃ NSCl.	1.	2.	Found. 3.	4.	5.
C	47.92	48.29	47.84			
H	2.45	2.97	3.04			
N	4.30	4.32	4.42			
S	9.83	10.33	10.56	9.96	10.34	10.26
Cl	10.90	10.90	11.11	11.03		

The two following structural formulas are possible for a substance derived from the unsymmetrical chloride, and having the composition indicated by the analyses.



The fact that the product is apparently not acted upon by alcoholic potash, together with its high melting-point and its properties generally favor the belief that it has the structure represented by the first.

The formation of the same product by the action of benzene and aluminium chloride on the symmetrical chloride seems to indicate that the product derived from each chloride is paranitroorthobenzoylbenzenesulphone chloride. This view agrees with the results obtained by the action of benzene and

aluminium chloride on orthosulphobenzoic acid by Saunders¹ and McKee.²

It is clear from the analysis that but one of the chlorine atoms of the chloride is replaced by this reaction. All attempts to prepare the diphenol derivative or paranitroorthobenzoylbenzenesulphone were unsuccessful, although the conditions were varied widely, both as to temperature and the length of time which the heating was continued.

The conditions already described give the best yield and also the purest product.

By allowing the mixture to stand for a day with occasional heating nearly to the boiling-point of the benzene, similar good results are obtained.

By heating to the boiling-point of benzene, for three hours, using a return-condenser, the yield is decreased, and a large amount of a black, tarry matter obtained, which is almost insoluble in benzene. This dissolves readily in absolute alcohol, from which solution it is precipitated as a dark red-colored powder on adding water, and after being thrown out of solution in this way, it becomes less soluble in absolute alcohol.

It swells up on heating and gives an odor like that obtained on burning sulphonic acids. On burning off the organic portion a considerable amount of alumina remains. Hydrochloric acid dissolves the alumina, leaving the organic portion in the form of a black, tarry mass.

V. The Action of Hydrochloric Acid on Paranitroorthobenzoylbenzenesulphone Chloride.

a. The Action of Dilute Hydrochloric Acid (sp. gr. 1.12).—The action of dilute hydrochloric acid was determined by boiling the sulphone chloride in a flask, provided with a return-condenser, with an excess of the acid until it was all dissolved, which usually requires about six hours. The solution is then filtered and evaporated on a water-bath, and the heating continued until no odor of hydrochloric acid remains.

¹ Am. Chem. J., **17**, 355.

² Inaug. Diss. Johns Hopkins University, 1895.

The resulting acid is obtained in the form of a dark solid substance which dissolves readily in water and takes up water on standing in the air.

Preparation of the Barium Salt.—The barium salt was prepared by adding barium carbonate to a solution of the acid in water, filtering off the excess of barium carbonate, and evaporating the solution under a bell-jar by means of a current of dry air.

The solution cannot be concentrated safely by boiling, as it causes a decomposition of the salt.

The barium salt was obtained in the form of small, light-colored crystals, arranged in tufts.

In the following analyses of salts, the base, as well as the sulphur, is calculated on the basis of the anhydrous salt.

0.2039 gram lost 0.0119 gram at 180° C. and gave 0.0586 gram BaSO₄.

0.2892 gram lost 0.0179 gram at 180° C. and gave 0.0810 gram BaSO₄.

	Calculated for (C ₁₃ H ₈ O ₆ NS) ₂ Ba + 3H ₂ O.	I.	Found. 2.
H ₂ O	6.72	5.84	6.18
Ba	18.29	18.12	17.65

The barium salt of another portion of acid, prepared in the same way, was obtained in the form of short, thick monoclinic prisms, which seemed to be made up of a series of plates.

0.2010 gram lost 0.0260 gram at 210° C. and gave 0.0557 gram BaSO₄.

0.2021 gram lost 0.0250 gram at 210° C. and gave 0.0562 gram BaSO₄.

0.2263 gram gave 0.1262 gram BaSO₄.

0.2109 " " 0.1201 " "

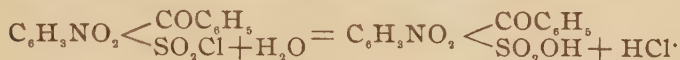
	Calculated for (C ₁₃ H ₈ O ₆ NS) ₂ Ba + 6H ₂ O.	I.	Found. 2.
H ₂ O	12.60	12.93	12.66
Ba	18.29	18.64	18.70
S	8.54	8.76	8.93

This salt became opaque on standing in a specimen tube for one month, due to the loss of water of crystallization.

0.2133 gram of the opaque salt lost 0.0139 gram at 210° C.

	Calculated for $(C_{13}H_9O_6NS)_2Ba + 3H_2O$.	Found.
H ₂ O	6.72	6.56

The results of the analysis indicate that paranitroorthobenzoylbenzenesulphone chloride is converted into paranitro-orthobenzoylbenzenesulphonic acid by boiling with dilute hydrochloric acid, according to the following equation:



b. The Action of Concentrated Hydrochloric Acid, (sp. gr. 1.17).—The action of concentrated hydrochloric acid was determined by heating the sulphone chloride with a large excess of acid in a sealed tube.

The tube was first heated for six hours in a water-bath, but, as no action seemed to take place, it was transferred to a Carius furnace and heated for six hours at a temperature of 175° C. The substance dissolved, with the exception of a few dark flakes, and the acid was colored brown. The flakes were removed by filtration, the acid solution evaporated to dryness on a water-bath, and the heating continued until the resulting acid had no odor of hydrochloric acid. The barium salt was prepared as in the case of the acid derived from the sulphone chloride by the action of dilute hydrochloric acid. It crystallized in the form of light-colored, fine needles, which were arranged in loose tufts or clusters. A few darker crystals in the form of larger monoclinic crystals with rhombohedral habit were obtained from the mother liquor.

0.2103 gram of the needles lost 0.0164 gram at 210° C. and gave 0.0609 gram BaSO₄.

0.2083 gram of the needles lost 0.0158 gram at 210° C. and gave 0.0595 gram BaSO₄.

	Calculated for $(C_{13}H_9O_6NS)_2Ba + 3\frac{1}{2}H_2O$.	1.	Found.	2.
H ₂ O	7.75	7.69		7.59
Ba	18.29	18.21		18.20

0.1053 gram of the larger, dark crystals lost 0.0152 gram at 210° C. and gave 0.0279 gram BaSO_4 .

	Calculated for $(\text{C}_{13}\text{H}_8\text{O}_6\text{NS})_2\text{Ba} + 7\text{H}_2\text{O}$.	Found.
H_2O	14.40	14.43
Ba	18.29	18.20

VI. The Action of Dilute Sulphuric Acid on Paranitroorthobenzoylbenzenesulphone Chloride.

The action of sulphuric acid on the sulphone chloride was determined by heating in a flask with a return-condenser until it dissolved. The resulting product was an acid, which was converted into the barium salt by adding an excess of barium carbonate, as in the previous experiments.

The barium salt was obtained in the form of short needles arranged in clusters.

0.1943 gram lost 0.0119 gram at 180° C. and gave 0.0570 gram BaSO_4 .

	Calculated for $(\text{C}_{13}\text{H}_8\text{O}_6\text{NS})_2\text{Ba} + 3\text{H}_2\text{O}$.	Found.
H_2O	6.72	6.12
Ba	18.29	18.48

VII. The Action of Water on Paranitroorthobenzoylbenzenesulphone Chloride.

The action was determined by boiling the sulphone chloride in a flask with a return-condenser until it dissolved. It was necessary to boil somewhat longer to dissolve the substance than in the experiments in which acids were used.

The resulting product was an acid which, by treating in the usual way with barium carbonate, gave a barium salt which crystallized in well formed monoclinic crystals.

0.1610 gram lost 0.0207 gram at 210° C. and gave 0.0443 gram BaSO_4 .

	Calculated for $(\text{C}_{13}\text{H}_8\text{O}_6\text{NS})_2\text{Ba} + 6\text{H}_2\text{O}$.	Found.
H_2O	12.60	12.85
Ba	18.29	18.56

VIII. The Action of Absolute Alcohol on Paranitroorthobenzoylbenzenesulphone Chloride.

The action of absolute alcohol on the sulphone chloride

was determined by boiling in a flask with a return-condenser until it dissolved. It dissolved rather more rapidly than in the experiments in which acid was used, and the boiling was continued for a short time after all the material was dissolved. After a part of the alcohol was evaporated, a few drops of the solution showed indications of crystallization on evaporating rapidly on a watch-glass, but, on further evaporation, the solution darkened and the resulting product was an acid as in the previous experiments.

The barium salt was prepared as in the previous experiments. The crystals first obtained were in the form of small light-colored needles arranged in clusters, but later well formed monoclinic crystals were obtained from the same solution.

0.1957 gram of the needle-shaped crystals lost 0.0153 gram at 210° C. and gave 0.0559 gram BaSO_4 .

	Calculated for $(\text{C}_{13}\text{H}_9\text{O}_6\text{NS})_2\text{Ba} + 3\frac{1}{2}\text{H}_2\text{O}$.	Found.
H_2O	7.75	7.81
Ba	18.29	18.21

0.2007 gram of the larger crystals lost 0.0285 gram at 210° C. and gave 0.0535 gram BaSO_4 .

	Calculated for $(\text{C}_{13}\text{H}_9\text{O}_6\text{NS})_2\text{Ba} + 7\text{H}_2\text{O}$.	Found.
H_2O	14.40	14.26
Ba	18.29	18.27

The above analyses indicate that the action of dilute or concentrated hydrochloric acid, sulphuric acid, water and alcohol on paranitroorthobenzoylbenzenesulphone chloride converts it into paronitroorthobenzoylbenzenesulphonic acid.

IX. Comparison of the Barium Salts.

The analyses of the barium salt of the acids, derived from the action of the various substances on the sulphone chloride, show that the acid is in every case the same and that the salts contain the same amount of barium when calculated upon the basis of the anhydrous salt. The amount of water of crystallization varies widely in the different salts, depending on the conditions under which crystallization takes place.

The needles are obtained from the more concentrated solutions, and crystals of this form are first obtained from a solution which has been evaporated by heating before placing it under a bell-jar. All crystals having this form contain three or three and a half molecules of water of crystallization.

The larger monoclinic crystals which form in the same solution after the formation of needles ceases, or when a cold solution is evaporated to the point of crystallization under a bell-jar, are characterized by six molecules of water of crystallization. On exposure to the air or even in a stoppered tube these lose water of crystallization and become opaque.

The only analysis made of a crystal that had changed in this way shows that it contains three molecules, while it crystallized with six.

Those crystals which contain seven molecules of water of crystallization are obtained on slow crystallization, on standing in the air, form a dilute solution or form a mother-liquor from which crystals containing a less amount of water of crystallization have been deposited.

Although all the barium salts containing different amounts of water of crystallization appear to crystallize in the monoclinic system, they show clearly a variation in form.

Owing to lack of time, no comparative study could be made of the relation existing between the amount of water of crystallization and the crystallographic constants.

The barium salt of paranitroorthobenzoylbenzenesulphonic acid is characterized by an intense bitter taste.

X. Preparation of Other Salts from the Barium Salt of Paranitroorthobenzoylbenzenesulphonic Acid.

These were prepared from an aqueous solution of the barium salt by precipitating the barium exactly by means of sulphuric acid and neutralizing the free acid exactly with the carbonate of the base.

The solutions were evaporated to crystallization under a bell-jar by means of a current of dry air.

In the analyses the amount of the base is calculated on the basis of the anhydrous salt.

a. The Sodium Salt.—The sodium salt was obtained in the form of fine white crystals composed apparently of monoclinic prisms.

They appear to undergo no change on exposure to the air.

0.1898 gram lost 0.0105 gram at 210° C. and gave 0.0384 gram Na_2SO_4 .

0.2022 gram lost 0.0111 gram at 210° C. and gave 0.0402 gram Na_2SO_4 .

	Calculated for ($\text{C}_{13}\text{H}_8\text{O}_6\text{NS}$)Na + H_2O .	I.	Found. 2.
H_2O	5.19	5.53	5.49
Na	6.99	6.93	6.82

b. The Potassium Salt.—The potassium salt was obtained in the form of fine white needles which were too small to indicate the form of crystallization. They became opaque on exposure to the air.

0.2118 gram lost 0.0014 gram at 210° C. and gave 0.0546 gram K_2SO_4 .

0.2041 gram lost 0.0013 gram at 210° C. and gave 0.0518 gram K_2SO_4 .

	Calculated for ($\text{C}_{13}\text{H}_8\text{O}_6\text{NS}$)K.	I.	Found. 2.
K	11.33	11.63	11.46

c. The Magnesium Salt.—The magnesium salt was obtained in the form of tabular monoclinic crystals having a marked pearly lustre. Some of the crystals measured nearly a centimeter in length. They appear to undergo no change on exposure to the air.

0.1940 gram lost 0.0408 gram at 210° C. and gave 0.0294 gram MgSO_4 .

0.1997 gram lost 0.0425 gram at 210° C. and gave 0.0303 gram MgSO_4 .

	Calculated for ($\text{C}_{13}\text{H}_8\text{O}_6\text{NS}$) $_2\text{Mg}$ + $9\frac{1}{2}\text{H}_2\text{O}$.	I.	Found. 2.
H_2O	21.17	21.03	21.28
Mg	3.83	3.88	3.90

d. The Calcium Salt.—The calcium salt was obtained in the form of thin pearly plates having no regular bounding planes.

They become opaque on exposure to the air and crumble to a white powder.

0.1373 gram lost 0.0096 gram at 210° C. and gave 0.0280 gram CaSO_4 .

0.1199 gram lost 0.0082 gram at 210° C. and gave 0.0242 gram CaSO_4 .

	Calculated for $(\text{C}_{13}\text{H}_8\text{O}_6\text{NS})_2\text{Ca} + 3\text{H}_2\text{O}$.	1.	Found. 2.
H_2O	7.65	6.99	6.88
Ca	6.13	6.44	6.39

e. The Lead Salt.—The lead salt was obtained in clusters of small tabular monoclinic crystals, which became opaque very slowly on exposure to the air.

0.2121 gram lost 0.0219 gram at 210° C. and gave 0.0711 gram PbSO_4 .

0.2039 gram lost 0.0213 gram at 210° C. and gave 0.0699 gram PbSO_4 .

0.1543 gram lost 0.0166 gram at 210° C. and gave 0.0509 gram PbSO_4 .

	Calculated for $(\text{C}_{13}\text{H}_8\text{O}_6\text{NS})_2\text{Pb} + 5\frac{1}{2}\text{H}_2\text{O}$.	1.	Found. 2.	3.
H_2O	10.78	10.32	10.44	10.75
Pb	25.25	25.43	25.95	25.23

The copper salt underwent decomposition on evaporation.

XI. The Action of Phosphorus Pentachloride on the Sodium Salt of Paranitroorthobenzoylbenzenesulphonic Acid.

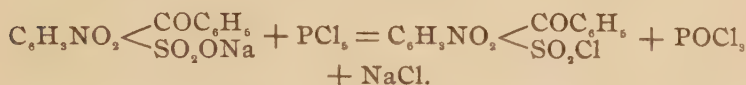
The sodium salt and phosphorus pentachloride in the proportion of 1 molecule to $1\frac{1}{2}$ were mixed by grinding together in an evaporating dish. There were no evidences of action, even upon adding a considerable quantity of phosphorus oxychloride, but on heating there was slight action.

The heating was continued for about ten minutes, and the pasty mass was then treated with a considerable volume of cold water. Most of the material dissolved, but part hardened to a solid mass. After carefully washing with water this material was washed with absolute alcohol, dissolved in benzene, and crystallized out by adding an equal volume of anhydrous ether. The product separated out as clusters of small,

light-colored crystals, which melted at 174–176° C. (uncor.) and as a scale, around the sides of the beaker, which melted at 160–170° C. (uncor.). It was entirely free from the dark purple material obtained as an impurity in the preparation of paranitroorthobenzoylbenzenesulphone chloride by the action of benzene and aluminium chloride.

A considerable portion of the material was insoluble in benzene and melted at 240–245° C. (uncor.).

The method of formation of this material, together with its melting-point, its solubility in benzene, from which it crystallizes readily upon the addition of absolute ether, indicate that it is paranitroorthobenzoylbenzenesulphone chloride. The method of formation from the sodium salt is indicated by the following equation :



The material melting at 174–176° C. was boiled in a flask with a return-condenser, with an excess of dilute hydrochloric acid until it was completely dissolved. This required seven hours. The solution was filtered and evaporated to dryness on a water-bath, and the heating continued until all hydrochloric acid was driven off. The product was a dark solid, similar to that obtained by the action of acid on paranitroorthobenzoylbenzenesulphone chloride. An excess of barium carbonate was added to an aqueous solution of the product, the excess of carbonate filtered off, and the solution, which had the characteristic bitter taste of the barium salt of paranitroorthobenzoylbenzenesulphonic acid, evaporated under a bell-jar. On evaporation the solution yielded a small amount of a crystalline product containing barium, which shows that the product is the barium salt of an acid.

The formation of paranitroorthobenzoylbenzenesulphonic acid by the action of hydrochloric acid on the product of the action of phosphorus pentachloride on the sodium salt of paranitroorthobenzoylbenzenesulphonic acid, and its conversion into the barium salt confirms the view already expressed that

the action of phosphorus pentachloride on the sodium salt gives the sulphone chloride.

XII. The Action of Concentrated Ammonia on Paranitroorthobenzoylbenzenesulphone Chloride.

a. The Action of Concentrated Ammonia on the Chloride in an Open Vessel.—Some of the sulphone chloride boiled with concentrated ammonia in a test-tube was apparently but little changed, but, on filtering off the unchanged chloride and evaporating the ammonia, a yellow amorphous powder was obtained which melted at 234°C . (uncor.) after washing with water to remove any ammonium chloride.

The action of ammonia on the chloride was next tried by boiling in a flask, provided with a return-condenser, to which fresh portions of ammonia were added as the strength decreased by evaporation. After boiling in this way for about three hours the ammonia became yellow, although the material remaining undissolved was not changed in appearance. It was found, however, that the insoluble material melted at 234°C . (uncor.) and that it was unchanged by crystallization from benzene and alcohol. A small amount of material was also obtained which melted above 375°C . (uncor.). This indicated that the chloride had undergone a change on heating with ammonia.

b. The Action of Concentrated Ammonia on the Chloride in a Sealed Tube.—In the experiments in which the chloride was heated with concentrated ammonia in an open vessel, the strength of the ammonia decreased, and, even though continually replaced, the experiment was complicated by the action of water. In order to avoid this, the chloride was heated with concentrated ammonia in a sealed tube. Upon heating it as long as it had been found necessary to heat in an open vessel in order to effect the transformation, the product was mixed with a considerable amount of red-colored material which melted above 275°C . (uncor.).

As the result of several experiments it was found that, by heating the chloride in a sealed tube for two or two and a half

hours in a water-bath, it is mainly converted into a clear, granular product which melts at 234°C . (uncor.) as in the previous experiment.

The small amount of the dark high-melting product formed exists as a thin coating and can easily be removed mechanically or dissolved in alcohol, which dissolves it readily without dissolving the main product.

The material thus prepared is obtained in the form of a light-green, granular powder, having a constant melting-point of 234°C . (uncor.). The substance contains no chlorine.

0.2014 gram gave 15.81 cc. N.

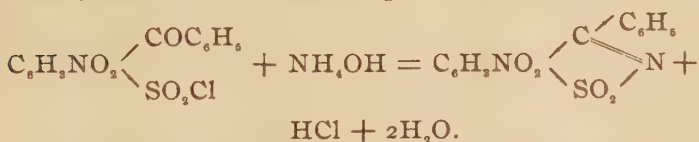
0.1970 gram gave 15.67 cc. N.

0.2075 gram gave 0.1756 gram BaSO_4 .

0.2011 gram gave 0.1692 gram BaSO_4 .

	Calculated for	Found.	
	$\text{C}_6\text{H}_5\text{NO}_2 \begin{array}{l} \diagup \text{C} \diagdown \\ \diagdown \text{SO}_2 \diagup \end{array} \begin{array}{l} \diagup \text{C}_6\text{H}_5 \\ \diagdown \text{N} \end{array}$	I.	2.
N	9.72	9.86	9.99
S	11.11	11.60	11.55

The results of analysis, together with those described in the following section, indicate that the main product of the action of concentrated ammonia on the chloride is the lactim of paranitroorthobenzoylbenzenesulphonic acid. The reaction by which it is formed is represented as follows :



The lactim is insoluble in water, only slightly soluble in alcohol and readily soluble in benzene.

The formation of the lactim of the sulphonic acid by the action of concentrated ammonia agrees with the formation of the lactim of orthobenzoylbenzenesulphonic acid by the action of dry ammonia gas on the sulphone chloride as observed by Saunders.¹

¹ Am. Chem. J., 17, 359.

XIII. The Action of Concentrated Ammonia on the Lactim of Paranitroorthobenzoylbenzenesulphonic Acid.

The presence of a red-colored amorphous product, melting above 275° C. (uncor.), with the lactim formed by the action of ammonia on the chloride, together with the fact that the amount of this product was increased as the length of time of heating was increased, indicated that another product was formed by the continued action of ammonia. A considerable quantity of this material was prepared by heating some of the paranitroorthobenzoylbenzenesulphone chloride in a sealed tube until the only product consisted of the red-colored substance desired. It was found necessary to heat it to the temperature of the water-bath for twenty-four hours in order to effect this transformation, while two and a half hours were sufficient to transform the sulphone chloride into the lactim.

The product is insoluble in water, but dissolves readily in absolute alcohol, giving a red solution with a marked green fluorescence. It is thrown out of solution by adding a considerable volume of water.

On evaporating the alcoholic solution it is deposited as a red-colored crust which seems to possess no crystalline structure.

0.2150 gram gave 19.24 cc. $N = 11.24$ per cent. N.

0.1615 gram gave 0.1200 gram $BaSO_4 = 10.20$ per cent. S.

The results of the analysis show that while the percentage of sulphur remains about the same as in the lactim, the percentage of nitrogen is increased, but not to an amount corresponding to the composition of any substance likely to be derived from the lactim by the further action of ammonia.

These results, together with the impossibility of obtaining the product in crystalline condition and its properties generally, indicate that it is probably not a definite chemical compound, and that the lactim probably undergoes decomposition by the further action of ammonia.

XIV. The Action of Dilute Hydrochloric Acid on the Lactim of Paranitroorthobenzoylbenzenesulphonic Acid.

The action of hydrochloric acid on the lactim was first tried

by boiling in a flask, provided with a return-condenser, with an excess of acid. The lactim showed but little change after boiling with the acid for thirty hours. The acid was colored yellow, but this was found to be due to solution of the lactim. By evaporating off the acid, the lactim is recovered with its melting-point unchanged.

By heating the lactim with a large excess of hydrochloric acid in a closed tube to 150° – 175° C. in a furnace for five hours, about half of the lactim is dissolved and is not deposited on cooling. On heating to 200° C. for seven hours longer, all of the lactim goes into solution, and is not deposited on cooling, and the acid has a dark yellow color. On evaporating the filtered acid solution, a yellow, crystalline product is obtained, which has not a constant melting-point. The melting-point, immediately after pressing out between filter-paper, is 100° – 160° C. (uncor.), and it is charred by heating to 210° C. (uncor.) in an air-bath. If, however, it is first carefully dried, it appears to melt at a much higher temperature.

This indicates that the product is a salt which melts in its water of crystallization.

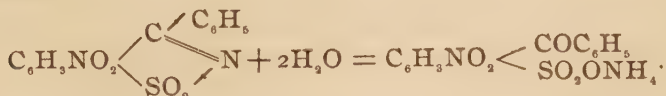
Analysis of a sample carefully dried :

0.2009 gram gave 14.33 cc. N.

0.1692 gram gave 0.1264 gram BaSO_4 .

	Calculated for $\text{C}_6\text{H}_3\text{NO}_2 \begin{smallmatrix} \text{COC}_6\text{H}_5 \\ \text{SO}_2\text{ONH}_4 \end{smallmatrix}$	Found.
N	8.64	8.95
S	9.88	10.24

The results of analysis indicate that the product is the ammonium salt of paranitroorthobenzoylbenzenesulphonic acid. The transformation takes place according to the following equation :



The ammonium salt thus obtained has generally the form of a yellow crystalline powder, but under the conditions existing in one of the experiments, a few thick, needle-shaped

crystals about a centimeter long were obtained. It dissolves readily in water.

XV. Conclusion.

The principal results obtained in the foregoing investigation may be briefly stated as follows :

By using phosphorus pentachloride in the proportion of two and a half molecules to one of the anhydrous acid potassium salt and heating for five hours under the conditions indicated, the unsymmetrical chloride is the only product.

This may be crystallized readily in any quantity at the temperature of the laboratory by using ligroin (90° – 125°) as the solvent, provided the impurities are first removed by washing with ligroin (50° – 80°).

The action of benzene and aluminum chloride on the symmetrical and on the unsymmetrical chloride gives, in both cases, paranitroorthobenzoylbenzenesulphone chloride.

The action of hydrochloric acid, concentrated or dilute, dilute sulphuric acid, water and alcohol on paranitroorthobenzoylbenzenesulphone chloride is the same. The product formed is, in each case, paranitroorthobenzoylbenzenesulphonic acid, which forms a series of characteristic salts.

The action of phosphorus pentachloride on the sodium salt of paranitroorthobenzoylbenzenesulphonic acid gives rise to the formation of paranitroorthobenzoylbenzenesulphone chloride identical with that from which the acid was derived by the action of acids or water.

The action of concentrated ammonia on paranitroorthobenzoylbenzenesulphone chloride for a limited length of time gives the lactim of paranitroorthobenzoylbenzenesulphonic acid.

The further action of concentrated ammonia gives a substance of indefinite composition, which probably indicates a decomposition of the lactim, which is first formed.

The continued action of concentrated hydrochloric acid at a high temperature in a sealed tube converts the lactim into the ammonium salt of paranitroorthobenzoylbenzenesulphonic acid.

BIOGRAPHICAL.

The author was born at Dedham, Mass., August 26, 1867. His early education was received in the public schools of Newton, Mass. In 1885 he entered the Massachusetts Institute of Technology and received the degree of Bachelor of Science in 1890. From 1890 to 1893 he was engaged in experimental work in connection with the water supply of Boston, Mass. In 1893 he entered the Johns Hopkins University, where he has since pursued courses in chemistry, mineralogy and geology.

